

# **Evaluating the Health Outcomes from Newborn Screening for Cystic Fibrosis**

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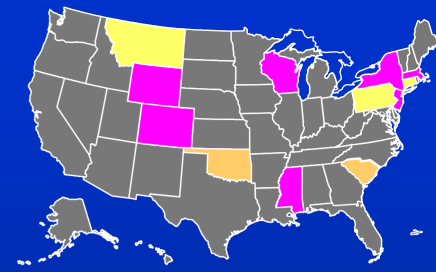
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Centers for Disease Control and Prevention

National Center on Birth Defects and Developmental Disabilities



## Public Health Context



- About 20% of U.S. infants are screened for cystic fibrosis (CF)
- State by state decision, mostly since 1999
- No national public health consensus
- CDC workshops
  - 1997 insufficient evidence for routine screening
  - 2003 evidence of moderate benefit



### **Time Line**

- January 1997 workshop convened by CDC & partners
- December 1997 MMWR Reports & Recommendations
  - Encourage pilot screening and research
  - Collect evidence on additional outcomes
  - Convene panel in 2 years to review new evidence
- May 2002 CF Foundation proposes new workshop
- January 2003 CDC/NCBDDD considers workshop
- April 2003 Experts visit CDC to present evidence
- November 2003 Workshop convened by CDC & CFF
- October 2004 MMWR Reports & Recommendations



## Age of Diagnosis in United States

- About 25% of children with CF are diagnosed soon after birth in absence of NBS
  - Meconium ileus
  - Prenatal diagnosis, family history, etc.
- Median age of diagnosis for others is 14 months
- With newborn screening (NBS), diagnosis is feasible within 1-2 months, about 12 months sooner



## **Arguments for Screening Infants for CF**

- Clinical utility improved outcomes
- Prevent diagnostic odyssey
- Opportunity for early treatment
  - Pancreatic enzymes
  - Vitamin supplements
  - High-fat dietary regimen
  - More aggressive antibiotic therapy
- Genetic counseling 1 in 4 risk of recurrence in siblings



### **Traditional Criteria for NBS**

- Clinical utility
  - Prevention of child death or severe disability
  - Model is PKU
- Other criteria
  - Frequency of condition
  - Feasibility and accuracy of screening test in DBS
  - Availability of treatment
  - Cost of screening, etc.
- No consideration of other benefits
  - Reduced morbidity
  - Improved quality of life
  - Benefits to families



## **Assessing Health Outcomes for CF**

- Traditional NBS criteria too narrow
  - CF not associated with intellectual disability
  - Child deaths not common in CF
- Direct clinical outcomes
  - Malnutrition and growth retardation
  - Lung disease
- Indirect outcomes
  - Cognitive development
  - Health-related quality of life (HRQoL)
  - Hospitalizations and burden of treatment
- Balance of outcomes
  - Risks and benefits
  - Cost-effectiveness



### Which Health Outcomes Matter Most?

- Those of direct concern to patients and families
- Strength of Recommendation Taxonomy (SORT) (Ebell et al. 2004):
  - Disease-oriented outcomes

"intermediate, histopathologic, physiologic, or surrogate results...that may or may not reflect improvements in patient outcomes"

Patient-oriented outcomes

"matter directly to patients and help them live longer or better lives, including reduced morbidity, reduced mortality, symptom improvement, improved quality of life, or lower cost"



## Classifying Outcomes in CF

- Disease-oriented outcomes intermediate outcomes measured in routine CF care
  - Growth parameters
  - Lung function and x-rays
- Patient-oriented outcomes
  - Survival
  - Cognitive function
  - Health-related quality of life
  - Hospitalizations, intensive therapies, costs
- Matter of degree
  - Large decrements of direct concern to families
  - Example: growth hormone therapy



## **Assessing Outcomes at 1997 Workshop**

- **Evidence from three studies** 
  - Wisconsin RCT, children born 1985-1994
  - Australia observational study with historical controls, children born 1978-1984
  - Netherlands observational study with nonrandomized controls, children born 1973-1979

#### Conclusions

- Potential biases in both observational studies
- Consistent evidence of nutritional outcomes
  - Improved height-for-age
  - Reduced growth retardation (below 5<sup>th</sup> centile)
- No agreement of sufficient basis for routine NBS
- Need for evidence on other outcomes (cognitive, **HRQoL**, cost-effectiveness)



## Challenges in Interpreting Evidence: Limitations of Individual Studies

- Biases in observational and some clinical studies
  - Ascertainment bias
  - Differences in genotypes, ethnicity, etc.
  - Differences in care provided
- Randomized controlled trials
  - Chance differences between groups
  - Other threats to validity contamination
- Common issues
  - Adequate follow-up time and loss to attrition
  - Statistical power number of observations



## **Example: Wisconsin RCT**

- Well-designed trial all children screened 1985-1994, randomized to early diagnosis or blinding, 18 year follow-up
- Small numbers of children
  - Screened w/o MI (n=56)
  - Controls w/o MI (n=48)
- Chance difference between groups
  - ΔF508 homozygotes more common in screened group, 59% vs 47% (p<0.001)
- Contamination of pulmonary outcomes
  - One center exposed infants to older, infected patients
    - Median age of colonization with Ps. aeruginosa
      - 1.0 years for screened children at that center
      - 4.5 years for control children at same center
      - 5.6 years for screened children at other center
  - Poorer pulmonary outcome greater deterioration of chest x-rays with increasing age in screened group



## **Example: UK RCT**

- Children in Wales and West Midlands randomized to be screened or not, 1985-89
  - Screened (n=58)
  - Not screened (n=44)
- Limitations
  - No standardized treatment protocol
  - Incomplete ascertainment in unscreened cohort
  - Short follow-up: n=19 followed for 4 years
- Outcomes
  - No differences at 4 years (Chatfield et al. 1991)
  - Survival to 5 years (Doull et al. 2001)
    - Ascertained deaths from multiple sources
    - 4 CF-related deaths in unscreened cohort
    - 0 CF-related deaths in screened cohort
    - Difference is significant (p<0.05)</li>



# Challenges in Interpreting Evidence: Synthesizing Findings

- Statistical significance
  - Individual studies may be under-powered
  - Look for consistency of size of effect
- Assessing bias
  - Are reported prevalences in screened and unscreened groups comparable?
  - Are treatment protocols similar?
  - Is distribution of genotypes similar?
- Inconsistent findings
  - If outcomes depend on treatments provided, no consistent impact of screening may be expected
  - Consider exceptional factors in studies with discrepant findings (e.g. Wisconsin RCT)



## **Example: French Observational Study** (Siret et al., 2003)

- 1989-98 birth cohorts in neighboring regions, excluding children with meconium ileus
  - Brittany, with NBS for CF (n=77)
  - Loire-Atlantique, no NBS for CF (n=36)
- Comparability
  - Same birth prevalence of CF
  - Same treatment protocols
  - ΔF508 homozygotes more common in Brittany (not significantly) different)
- Outcomes for Brittany vs Loire-Atlantique
  - CF-related deaths 0/77 vs 3/36 (p<0.05)</li>
  - Hospitalization 49% vs 86% (p<0.0001)</li>
  - Height-for-age Z-scores 0.3-0.6 higher at 1, 3, 5 years (p<0.05)</li>
  - Better chest x-ray and clinical scores (p<0.05)
- Consistency: all findings consistent with other studies



# Cognitive Outcome in WI RCT: Overall (Koscik et al., 2004)

- Background
  - Malnutrition can affect neurodevelopment
  - Head circumference-for-age lower at diagnosis in control group
- Cognitive assessments
  - Conducted at ages 7-18 years (n=89)
  - Test of Cognitive Skills, 2<sup>nd</sup> Edition
  - CSI scale (similar to IQ)
- Findings for children without MI (n=71)
  - CSI Mean (SD) not significant (P=0.24)
    - Screened 104.6 (14.4)
    - Controls 99.8 (18.5)
  - No significant correlation of CSI with head circumference (P=0.11)



## Cognitive Outcome in WI RCT: Vitamin E (Koscik et al., 2004)

- Fat-soluble vitamin deficiencies common in CF
  - Vitamins A, E, and K
  - About half of children in WI RCT had plasma alphatocopherol < 300E at diagnosis – vitamin E deficiency
  - Deficiency corrected by vitamin supplements
- Findings of cognitive assessments (n=66)
  - CSI difference of 12.5 points (P<0.05) between Screen and Control children with early vitamin E deficiency

	Control,	Control,	Screen α-	Screen,
	α-T<300	α-T≥300	T<300	α-T≥300
	(n=16)	(n=13)	(n=17)	(n=20)
CSI Mean	91.5	107.7	104.0	105.8
(SD)	(15.1)	(15.4)	(16.2)	(15.0)



## Summary of Evidence on **Health Outcomes**

- Moderate impact on growth 0.3 Z-score difference in height-for-age
- Moderate impact on cognition overall difference of 5-6 IQ points in WI study
- Reduction in CF-related child mortality is reported in studies at ~50% or more
- Reduction in hospitalization and cost is possible
- No consistent improvement in pulmonary outcomes and some risk of harm without adequate infection control

